

Metzincin's canonical methionine is responsible for the structural integrity of the zinc-binding site

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Abstract

The metzincins constitute a subclan of metalloproteases possessing a HEXHHXXGXXH/D zinc-binding consensus sequence where the three histidines are zinc ligands and the glutamic acid is the catalytic base. A completely conserved methionine is located downstream of this motif. Families of the metzincin clan comprise, besides others, astacins, adamalysins proteases, matrix metalloproteases, and serralysins. The latter are extracellular 50 kDa proteases secreted by Gram-negative bacteria via a type I secretion system. While there is a large body of structural and biochemical information available, the function of the conserved methionine has not been convincingly clarified yet. Here, we present the crystal structures of a number of mutants of the serralysin member protease C with the conserved methionine being replaced by Ile, Ala, and His. Together with our former report on the leucine and cysteine mutants, we demonstrate here that replacement of the methionine side chain results in an increasing distortion of the zinc-binding geometry, especially pronounced in the χ^2 angles of the first and third histidine of the consensus sequence. This is correlated with an increasing loss of proteolytic activity and a sharp increase of flexibility of large segments of the polypeptide chain.

Keywords: crystal structure; directed mutagenesis; metalloprotease; metzincin; serralysin.

Introduction

Zinc-dependent endo-metalloproteases occur in all kingdoms of life and are involved in manifold biological functions. Many of them are characterized by a consensus sequence HEXXH, where the histidines are zinc ligands and the glutamic acid functions as catalytic base. A third zinc ligand is provided by an amino acid side-chain (His, Glu, Asp) downstream of this motif and the coordination

sphere is completed by the activated water molecule that attacks the peptide bond in the course of the reaction (Jiang and Bond, 1992; Tronrud et al., 1992). These enzymes are grouped in clan MA of the MEROPS protease database (Rawlings et al., 2006). The metzincins are a subclan, possessing an elongated HEXXH-XXGXX(H/D) zinc-binding motif with the last amino acid (mostly His) being the third zinc ligand (Bode et al., 1993; Stocker et al., 1995; Gomis-Ruth, 2003, 2009). The name of this subclan originates from a conserved methionine located in a 1,4-turn (so-called 'Met-turn'), which is sub-jacent to the base of the pyramidal arrangement of the three zinc ligands (Bode et al., 1993). Protease families belonging to the metzincins are, besides others, the astacins (Bond and Beynon, 1995), matrix metalloproteases (MMPs; Nagase et al., 2006), adamalysins (Blöbel, 2005), and serralysins (Maeda and Morihara, 1995). The latter are proteases secreted by Gram-negative bacteria utilizing a type I secretion system. Typical members are serralysin from *Serratia marcescens*, alkaline protease from *Pseudomonas aeruginosa* and protease C (PrTC) from *Erwinia chrysanthemi*. These 50 kDa proteins possess, besides an N-terminal approximately 220 amino acid long protease domain, a C-terminal β -roll moiety which is formed by tandem repeats of the glycine-rich nonapeptide GGXGXDXLX. This domain binds calcium and probably functions as an intramolecular chaperone (Baumann et al., 1993; Meier et al., 2007) (Figure 1A). Serralysins are classical metzincins where His-188, His-192, and His-198 are the zinc ligands and the Met-turn possesses the sequence 224-SIMSY-228. The tyrosine (Tyr-228) in this element stabilizes the tetrahedral transition state of the hydrolysis reaction (Hege and Baumann, 2001b) and is not present in all metzincin families (Figure 1B). Another unique feature of the serralysins is the loop comprising residues 201–210 in PrTC (drawn in orange in Figure 1), which is quite flexible in the *apo*-state and undergoes an induced-fit closure upon substrate binding.

The various metzincin families are rather distant to each other in evolution, exhibiting sequence identities of typically much less than 20%, and sequence alignments frequently pick up only the zinc-binding consensus motif. Between these families, the Met-turn is located in a non-conserved sequence stretch and the spacing between zinc-binding consensus motif and Met-turn varies. Nevertheless, as identified by structural alignment, the methionine always occupies the same spatial position and is in the same conformation (Figure 1). It lies below the base of the pyramid formed by the three histidines which are the amino acid zinc ligands. The sulfur atom is approximately 6 Å away from the metal ion and its lone pairs point in the opposite direction. The carbonyl oxygen accepts a hydrogen bond from the N δ 1 of His-188, which is the first zinc ligand.

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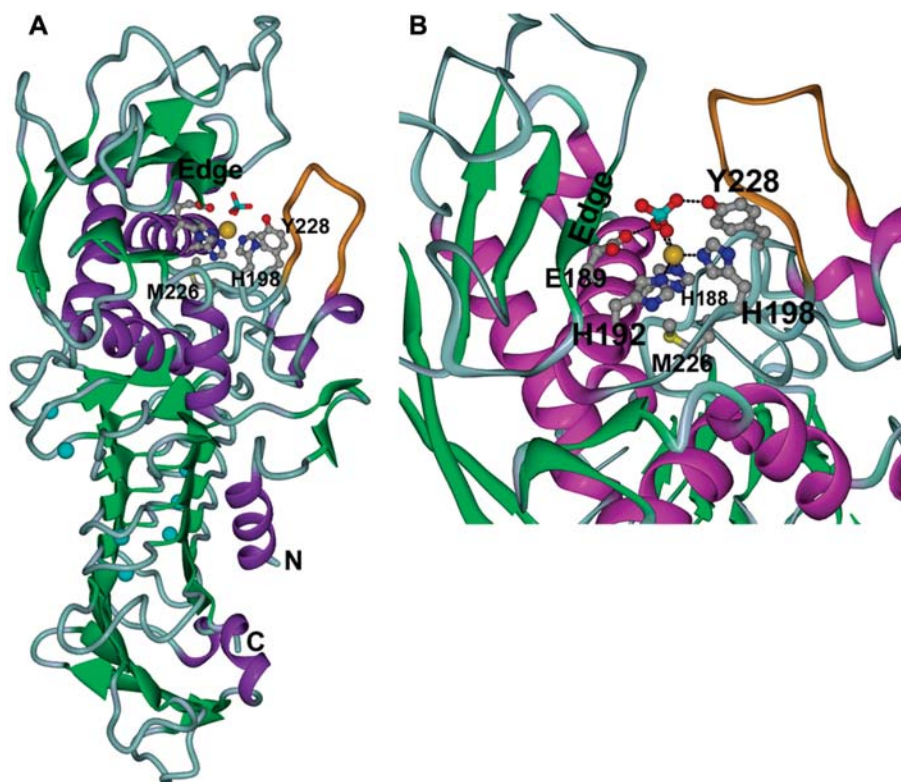


Figure 1 Overall structure and active center of wild-type PrtC.

(A) Ribbon diagram. The proteolytic domain is at the top, the β -roll domain at the bottom. Zinc is shown as golden sphere, calcium ions in light blue. A phosphate ion from the crystallization buffer coordinating to the zinc is shown as cyan and red sticks. The flexible loop (residues 200:206) is shown in orange. (B) Close-up of the active site. The substrate-binding edge strand is labeled as 'edge'. The sulfur atom of the methionine is depicted in yellow.

Methionine is not a conservative amino acid and ranks among the top 25% most rapidly mutating amino acids, being replaced most frequently by leucine, isoleucine, or valine (Jones et al., 1992; Tourasse and Li, 2000). Hence, owing to the apparently perfect conservation in a large set of very distantly related proteins, one should expect that the Met-turn methionine is very important for proteolytic activity and/or structural integrity. However, experimental studies replacing this residue by other amino acids in different metzincins did not deliver unequivocal results. In the case of MMPs, substitution of Met by selenomethionine (Qoronfleh et al., 1995; Pieper et al., 1997) or Leu or Ser (Butler et al., 2004) showed none or only minor changes in catalytic activity. Similar results were obtained by introduction of difluoromethionine in the serralsin alkaline protease (Walasek and Honek, 2005). On the other hand, mutation to Leu, Ile, or Ser in tumor necrosis factor α -converting enzyme led to a loss of ectodomain shedding (Perez et al., 2007). Our previous mutagenesis study of PrtC showed decreased proteolytic activity of mutants bearing Leu, Ile, and Ala at the position of Met-226. Crystal structures of the leucine mutant M226L and a double mutant EM189,226KC revealed subtle changes in the active site (Hege and Baumann, 2001a). The latter construct had an unwanted additional mutation of the active site glutamic acid, thus inactivating the enzyme completely. This allowed for more convenient purification and crystallization procedures.

In order to put the effects of methionine mutations on a more extended structural basis, we prepared more mutants and characterized their proteolytic activity. Fur-

thermore, we determined the crystal structures of double mutants carrying Ile, Ala, and His at the methionine position and report the results here.

Results

Proteolytic activity

In the past, we have reported Michaelis-Menten parameters for the M226L, M226I, and M226A mutants (Hege and Baumann, 2001a). Other mutants, such as M226H and M226N, could not be purified. Since the proteins were secreted into the extracellular medium and no other proteases were present there, we assayed proteolytic activities directly from the supernatant employing a resorufin-casein assay, normalized for protein concentrations determined by a BioRad assay as reported previously (Hege and Baumann, 2001b; see Figure 2). The results are in qualitative agreement with our previous data. Not surprisingly, the specific activity decreases Met>Leu>Ile>His>Ala, with the conservative mutations Leu and Ile having some 85% and 50% of the wild-type activity, respectively. The levels of secreted protein decrease in the same order, indicating some defect in synthesis and secretion or stability of the mutants.

Crystal structure analysis

Since yield and purity of some of the single methionine mutants was not sufficient for structural studies, we

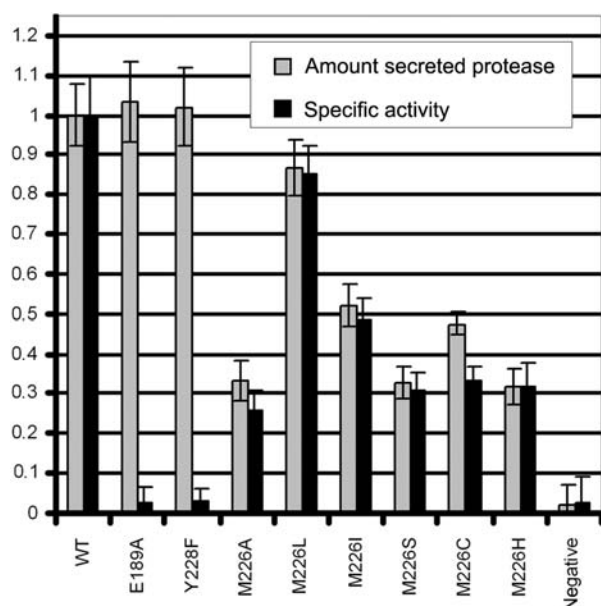


Figure 2 Activity assays.

Activity was assayed using resorufin-casein and media supernatant after spinning down the cells as described in the materials and methods section. Shown is the average of three independent measurements. Light bars indicate the ratio of secreted protein to cell density, and the dark bars indicate the specific activity. Values for the wild-type protein were normalized to 1.0. Cells carrying only the transport machinery plasmid were used as negative control ('negative').

introduced an additional mutation E189A, knocking out the catalytic base and thus inactivating the mutants completely. This greatly helped in purification and crys-

tallization. Relevant statistics are given in Table 1. No significant changes in the active site were introduced by this mutation (data not shown), besides the disappearance of a zinc-bound phosphate ion originating from the crystallization buffer and a slight movement of His-198. Therefore, we conclude that the differences we observe for the Met-226 mutants originate largely from the replaced methionine (Qoronfle et al., 1995; Pieper et al., 1997; Walasek and Honek, 2005).

An overlay of the active sites of the more conservative mutants (M226L, M226I, M226C) shows mainly a change in the geometry of the imidazole side chains of the coordinating histidine residues (Figure 3). In particular, the χ^1 angle of His-188 increases with decreasing activity (Table 2) and with decreasing size of the side-chain at position 226. This change in side-chain conformation causes a movement of the imidazole ring towards the cavity created by the mutation. The effect is most dramatic by cutting down the methionine side-chain to the C^β atom, i.e., in the M226A mutant (Figure 4). As can be seen in Figure 4A, the zinc, which is clearly present as documented by a large peak in the anomalous difference Fourier map, moves downwards by approximately 3 Å, while the imidazole side-chain of His-188 is nearly perpendicular to the orientation found in the wild-type enzyme. In order to keep the coordination sphere, His-198 changes position as well. This latter change is possible due to a dramatic order-disorder transition: amino acids 18–61 and 199–238, all very well defined in all the other mutants with the exception of M226H, become completely disordered. The second segment comprises the Met-turn and would clash with the new side-chain conformation of His-198. In other words, the polypeptide chain cov-

Table 1 Data collection and refinement.

Protein	M226I-E189A	M226A-E189A_inh	M226A-E189A_SLS	M226A-E189A_DESY	M226H-E189A
X-ray source	DESY X11	Ru300	SLS PXI	DESY X11	DESY X11
Unit cell a (Å) ^a	102.41	101.96	102.26	101.89	102.11
Unit cell c (Å)	121.03	123.22	123.40	122.52	122.86
Wavelength (Å)	0.812	1.542	1.281	0.812	0.812
Resolution (Å)	50.0–1.75	50.0–1.95	26.6–2.05	50.0–2.13	50.0–1.77
(outer shell)	(1.78–1.75)	(1.98–1.95)	(2.16–2.05)	(2.17–2.13)	(1.80–1.77)
R _{merge} (%)	4.8 (32.9)	3.4 (18.4)	11.7 (19.1)	8.4 (39.7)	4.0 (17.4)
Completeness (%)	99.9 (98.5)	99.6 (100.0)	99.9 (99.8)	99.9 (100.0)	100.0 (100.)
Redundancy	5.6 (3.8)	6.0 (5.1)	7.5 (5.8)	6.7 (3.8)	7.4 (5.7)
I/sigma (I)	23.0 (5.0)	28.3 (7.5)	15.5 (6.1)	12.9 (3.3)	45.1 (10.2)
refinement					
No. of reflections	74 320 (2183)	54 271 (1631)	47 147 (1412)	41 498 (1260)	72 453 (2138)
work (test) set					
No. atoms protein	3525	2937	2961	2947	3554
Ca ²⁺ /Zn ²⁺	7/1	7/1	7/1	7/0	7/2 ^c
Water	548	512	481	227	486
R/R _{free} (%)	15.5/17.9	15.3/16.6	14.8/17.0	18.2/20.7	16.6/18.7
RMS bonds (Å)	0.017	0.007	0.007	0.005	0.009
RMS angles (deg.)	1.56	1.0	0.97	0.88	1.63
Ramachandran	0	0	0	0	0
outliers (%) ^b					

^aCrystals belong to space group P3₁21.

^bAccording to Lovell et al. (2003) and Richardson (2003).

^cThe two Zn²⁺ ions occupy alternating sites with approximately 50% occupancy. The numbers in parentheses refer to the outer resolution shell.

ering the left side of the active center in Figure 4B–D (right side in Figure 1) becomes completely disordered. A patch of peptide-shaped electron density is clearly visible in the active site, forming an antiparallel β -strand with the edge of the sheet in the protease domain. It is unlikely that this peptide is one of the proteolysis products sometimes observed in the active centers of serrylsins (Baumann et al., 1993): firstly, the E189A mutation inactivates PrtC completely, and secondly the putative peptide extends over the zinc-binding position, which means it resembles more the educt than the products of the hydrolytic reaction.

However, the situation in the EM189,226AA mutant is even more complicated, owing to the additionally occurring complete loss of the zinc ion, as revealed by analysis of other crystals. Two out of four datasets show the condition depicted in Figure 4, while the remaining two crystals exhibit an entire loss of the zinc ion concomitant with a complete rearrangement of the active-site histidines (Figure 5, Table 2).

Another variation of the distortion of the active site could be observed in the EM189,226AH mutant. Here, an alternative zinc-binding site with approximately 50% occupancy, as derived by comparison of B-factors from neighboring atoms and the situation in other mutants, is created by the presence of the additional imidazole group, as is clearly revealed by the imaginary electron density component (Figure 6). His-188 occurs in two alternative conformations coordinating to both zinc sites. Polypeptide segments 18–61 and 199–238 exhibit slightly higher average B-factors than in the E189A

mutant, but are clearly visible with the exception of residues 203–210 belonging to the active center loop.

Discussion

Currently, some 800 members of the metzincin clan are known (Gomis-Ruth, 2003, 2009). All of them appear to possess the conserved methionine, although this residue is sometimes only tentatively identified by sequence alignments. Mutagenesis of such a conserved amino acid would be expected to exhibit large effects on activity and stability. However, conservative substitution by Leu or Ile, the naturally most frequently occurring mutations, displays only moderate effects. However, the previously reported distortions of the zinc environment have been confirmed, and they become increasingly larger with decreasing side-chain size at position 226. In order to assess the fit of the methionine side-chain compared to other amino acids, we calculated the shape complementarity value (SC value) (Lawrence and Colman, 1993) from our X-ray structures. A perfect fit would be indicated by a value of 1.0. The values for wild-type, M226L, M226I, and M226C mutants are 0.81, 0.73, 0.56, and 0.42, respectively. This clearly shows that in the jigsaw packing of hydrophobic amino acids at the base of the zinc-binding pyramid, methionine is best fitting. Thus, even replacement by leucine leads to observable distortions and a reduction in activity by approximately 15%. While this effect is small, it might nevertheless contribute to the selection pressure on the methionine. It should be noted

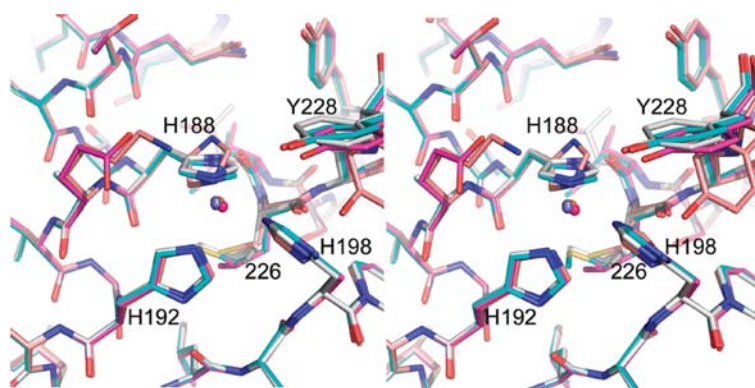


Figure 3 Active centers of M226 mutants revealing a distortion of the Zn^{2+} -binding site and a shift in the coordinating histidines. Stereo view of mutants E189A (cyan carbon atoms), M226L (magenta), EM189,226AI (gray), and EM189,226KC (wheat). The phosphate ion still present in the M226L structure has been omitted as well as the zinc-coordinating water molecules. The systematic change in the side-chain dihedrals, especially χ^2 of His-188, with decreasing size of the side-chain at position 226 is clearly visible (compare to Table 2).

Table 2 Side-chain dihedral angles of active-site histidines.

	Wild-type	E189A	M226L-E189A	M226I-E189A	M226C-E189K	M226H-E189A	M226A-E189A_inh	M226A-E189A_DESY
H188 χ^1	176	176	177	177	170	174	-169	-71
H188 χ^2	103	104	109	111	172	140	171	175
H192 χ^1	-73	-72	-68	-67	-67	-79	-71	-172
H192 χ^2	-44	-43	-44	-41	-38	-34	-41	104
H198 χ^1	-60	-68	-59	-61	-60	-69	-49	-49
H198 χ^2	145	152	143	143	125	126	111	112

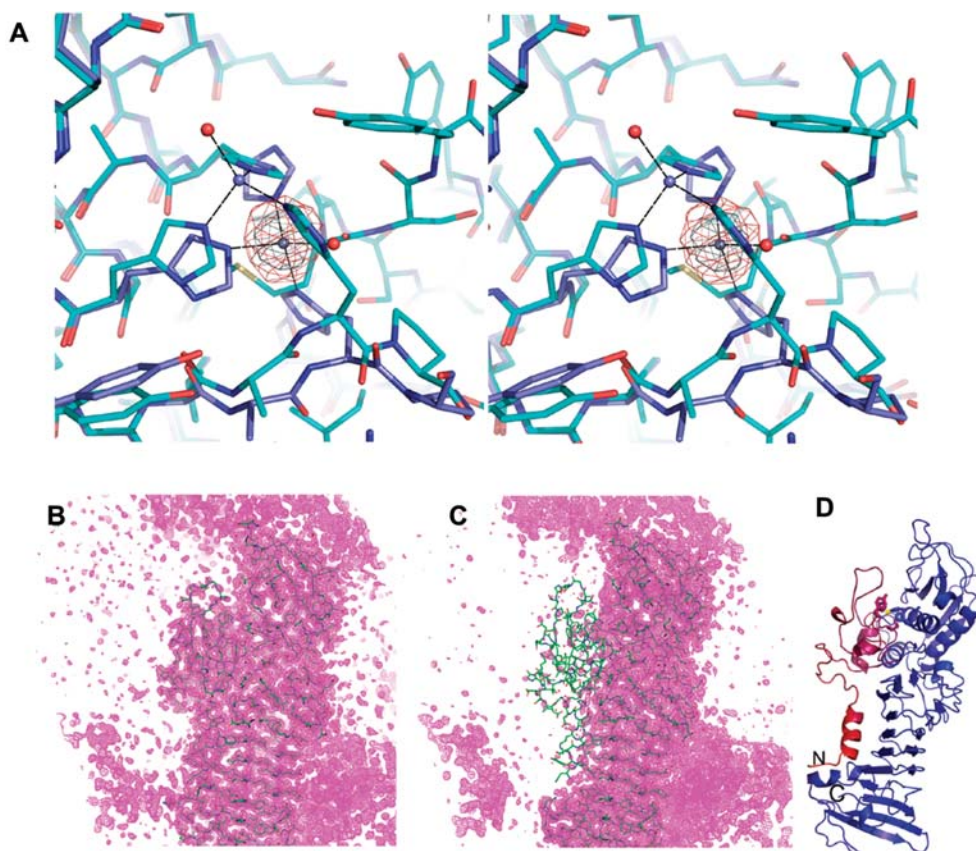


Figure 4 The EM189,226AA mutant as seen with the in-house and SLS data.

(A) Stereo view of an overlay E189A (cyan carbons, zinc shown as blue sphere) and EM189,226AA (blue carbon atoms, zinc depicted as gray sphere). Coordinating water molecules are shown as red spheres. An imaginary Fourier map is contoured at 10 and 15 standard deviations above the mean (magenta and gray mesh, respectively; data collected at SLS). (B) $2F_o - F_c$ electron density (contour level 1 sigma) of E189A. (C) $2F_o - F_c$ density map contoured at 1.0 sigma for the EM189,226AA double mutant. The orientation is similar as in panel (B). (D) Ribbon diagram of EM189,226AA colored according to B-factors from blue to red ($<25 \text{ \AA}^2$: blue; $>220 \text{ \AA}^2$: red). The orientation is approximately 180 degrees around the vertical from Figure 1 and similar as in panels (B) and (C). M226 and Y228 are shown as sticks as well as H188 and H192. The purple and red segments are completely invisible and have been omitted in panel (A).

that most other studies have replaced the methionine with selenomethionine or difluoromethionine, i.e., amino acids with different electronic properties at the δ -position of the side-chain but with a very similar size and shape

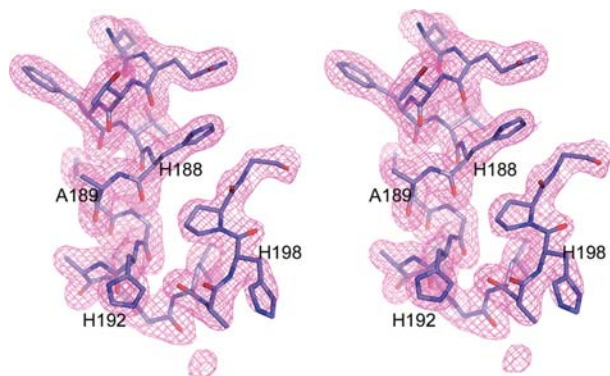


Figure 5 Active site of EM189,226AA as seen in data from DESY.

The $2F_o - F_c$ map (magenta) is contoured at 1.0 sigma above the mean. Anomalous difference Fourier maps show no signal for a Zn^{2+} ion anywhere close to the active site. This is in contrast to Figure 4A where the Zn^{2+} site is shifted.

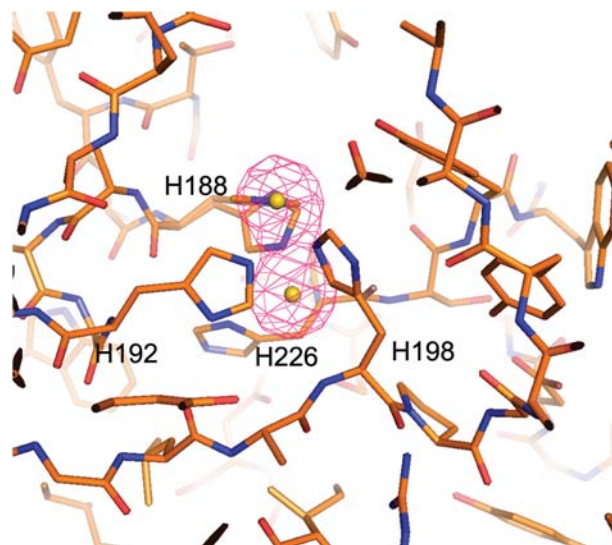


Figure 6 The EM189, 226AH mutant possesses two alternative Zn^{2+} sites.

The imaginary component of the electron density is shown in pink and contoured at 8 standard deviations above the mean (data collected at DESY). Zinc ions are shown as yellow spheres.

(Qoronfleh et al., 1995; Pieper et al., 1997; Walasek and Honek, 2005). Only the substitution of methionine by serine in MMP-2 (Butler et al., 2004) does not fit into our observations on PrtC. While in MMP2 no difference in activity and stability has been detected, in PrtC the M226S mutant exhibits approximately 30% of the wild-type activity, very similar to the cysteine mutant. Further studies will be needed to resolve these conflicting results and to attain a deeper understanding of the conservation of the methionine.

Materials and methods

Activity assay

LB medium (5 ml: 50 µg/ml kanamycin, 50 µg/ml chloramphenicol) was inoculated with a single colony carrying two plasmids pRUW520 (PrtC) and pRUW4inh1 (transport machinery) and incubated under shaking at 37°C overnight. The cell density was measured at 500 nm. Cells were removed by centrifugation and the protein concentration in the supernatant was measured by a Bradford assay. An aliquot was precipitated by trichloroacetic acid and analyzed on sodium dodecyl sulfate polyacrylamide gel electrophoresis. Proteolytic activity was assayed employing resorufin-casein (Roche Diagnostics, Basel, Switzerland) according to the manufacturer's instructions. Cells transformed only with pRUW4inh1 and grown in LB medium without kanamycin served as negative control.

Mutagenesis, protein expression, and purification

Site-directed mutagenesis was performed as described previously (Jones and Winistorfer, 1992; Hege and Baumann, 2001b) using the E189A DNA as template. Oligonucleotides were ordered from MWG Biotec (Ebersberg, Germany). The helper primer had the sequence 5'-CGC GGC CTC GAG CAA GAC GTT TCC CG-3'. The following mutagenic primers were employed (base changes are underlined):

M226I sense: 5'-CAA TTC AGT ATC ATA AGC TAC TGG GG-3';
M226A sense: 5'-CAA TTC AGT ATC GCG AGC TAC TGG GG-3';
and
M226H sense: 5'-CAA TTC AGT ATC CAT AGC TAC TGG GG-3'.

Expression was performed in *Escherichia coli* XL1 BLUE cells according to Delepelaire and Wandersman (1989). Briefly, cells were grown in LB medium containing 50 µg/ml kanamycin and 50 µg/ml chloramphenicol to an OD₆₀₀ of 0.6 at 37°C, then induced with 1 mM isopropyl-beta-D-thiogalactopyranoside (IPTG) and grown overnight at 37°C. Secreted proteins were purified by anion exchange chromatography as described previously (Hege and Baumann, 2001a,b). Trigonal-bipyramidal crystals grew in hanging drops at 20°C from a solution of 10 mg/ml protein and a reservoir containing 2 M NaCl, 0.1 M NaH₂PO₄, 0.1 M KH₂PO₄, 0.1 M 2-(N-morpholino)ethanesulfonic acid (MES), pH 6.5.

Data collection and structure refinement

Crystals were cryo-protected by the addition of 25% (v/v) glycerol and flash-cooled in a 110 K nitrogen stream. Data collection in-house employed a Rigaku (Tokyo, Japan) RU300 rotating anode and an RaxisIV image plate detector (Rigaku, Tokyo, Japan). Synchrotron data were collected at the EMBL outstation at Deutsches Elektronensynchrotron (DESY) in Hamburg (Germany) utilizing an MAR165 CCD detector. Data were processed

using the HKL2000 package (Otwinowski and Minor, 1997). Interactive model rebuilding was effected with O (Jones et al., 1991) and COOT (Emsley and Cowtan, 2004). Refinement was initially carried out using CNS (Brunger et al., 1998) and completed by PHENIX using TLS refinement (Adams et al., 2002). Figures were prepared with PYMOL (<http://www.pymol.org>). Coordinates of the various structures have been deposited in the Protein Data Bank (PDB) under the entry codes 3HB2, 3HBV, 3HBU, 3HBV, and 3HDA.

Acknowledgments

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